Two histological methods for recognition and study of cortical microinfarcts in thick sections

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Abstract

Cortical microinfarcts are the most widespread form of brain infarction but frequently remain undetected by standard neuroimaging protocols. Moreover, microinfarcts are only partially detectable in hematoxylin-eosin-stained 4-10 µm paraffin sections at routine neuropathological examination. In this short report, we provide two staining protocols for visualizing cortical microinfarcts in 100-300 µm sections. For low-power microscopy, the first protocol combines aldehyde fuchsin staining for detection of lipofuscin granules and macrophages with Darrow red counterstaining for Nissl material. The second protocol combines collagen IV immunohistochemistry with aldehyde fuchsin/Darrow red followed by enzymatic digestion (pepsin) to obtain successful collagen IV immunostaining (immuno-fluorescence) results of the microvasculature in formaldehyde-fixed human brain tissue; but the authors did not examine/apply this method to cortical microinfarcts. Clearly, both autopsy- and MRI-based studies will be needed, going forward, to determine the exact histopathology of these lesions.1,3

Materials and Methods

All procedures complied with the University of Ulm’s ethics committee guidelines and with German law governing usage of human post-mortem tissue. Two autopsy brains (2 males, 68 and 74 years of age) fixed for 14 days by immersion in a 4% aqueous solution of formaldehyde were used. Tissue blocks embedded in polyethylene glycol (PEG 1000, Merck, Carl Roth Ltd, Karlsruhe, Germany) were coronally sectioned from a tetramer (Jung, Heidelberg, Germany) at 100-300 µm as previously described.1,14 If paraffin-embedded tissue was preferred, the staining protocols provided below can be applied to sections of 70-100 µm thickness.13 Histologic sections were viewed and assessed with an Olympus BX61 microscope (Olympus Optical, Tokyo, Japan). Digital micrographs were taken with an Olympus XC50 camera using the Cell DÒ Imaging Software (Olympus, Münster, Germany). The program’s Extended Focal Imaging (EFI) function was used (Figure 1 d,f) to fuse stacks of four differently focused single images into one sharply focused image.

Protocol I - screening for cortical microinfarcts

To provide an overview of cortical microinfarcts for research purposes rather than for routine neuropathology in brain tissue blocks (Figure 2a), we used free-floating sections of 300 µm thickness. Whereas sections of this thickness are not sufficiently penetrated by all commercially available antibodies, they are ideal here for aldehyde fuchsin staining, which selectively visualizes intraneuronal and extraneuronal lipofuscin as well as CD68-positive macrophages in brain tissue. If desired, counterstaining for Nissl material with Darrow red can be performed (Figure 2b).15

1. Dissolve 1.5 g sodium stearate (Carl Roth, Karlsruhe, Germany) in 100 mL 70% ethanol to remove disruptive cerebrosides from the tissue. Rinse sections in sodium stearate for 1 h at 60°C. Rinse 5 min in 70% ethanol.
2. Dissolve 1 g sodium percarbonate
(Acros Organics, Geel, Belgium) in 100 mL 70% ethanol. Oxidize sections in sodium percarbonate for 1 h at room temperature after heating the sodium percarbonate solution at above 60°C. A white sediment, which is desirable, persists following dissolution. Agitate gently on a shaking table. Rinse sections 5 min in 70% ethanol.

3. Dissolve 5 mL aldehyde fuchsine solution (Morphisto, Frankfurt am Main, Germany) in 100 mL 70% ethanol and add 20 mL glacial acetic acid. Transfer specimens to aldehyde fuchsine solution for 1 h on a shaking table (gentle setting). Rinse sections twice in 70% ethanol to remove excess dye. When only aldehyde fuchsine staining is desirable, proceed to steps 5 and 6 immediately below.

4. Alternatively, transfer sections to deionized water and place into Darrow red solution for 1 h or more (agitate gently on a shaking table). To do so, dissolve 0.25 g Darrow red (Aldrich/Merck, Darmstadt, Germany) in 1000 mL deionized water, add 15 mL glacial acetic acid, boil for 20 min, filter solution, and add 250 mL of 0.2 Mol (i.e., 6.9 g sodium acetate trihydrate in 250 mL deionized water) sodium acetate solution (Merck, Darmstadt, Germany). No. Deionized water and place into Darrow red solution (30%) and 80 mL Tris. Rinse 5 min in Tris or PBS buffer. Agitate gently on a shaking table.

5. Dehydrate sections through a graded ethanol series. To obtain flatness, place free-floating sections between sheets of filter paper and compress gently between porcelain desiccator plates (140 mm diameter, Inv #119c, Morgan Technical Ceramics Haldenwanger, Germany, or Inv #8104, Lehman Scientific, Wrightsville, PA, USA) in 70%, 96% (30 min each), and 100% ethanol (30 min).

6. Remove sections from filter paper and dehydrate for another 30 min in 100% ethanol. A thorough dehydration step is crucial when using 300 µm sections. Clear in xylene twice (30 min each), coverslip in a high-refractive medium or synthetic resin (Histomount, National Diagnostics, Atlanta, GA, USA), and mount on slides.

Protocol II - for detailed study of cortical microinfarcts

For additional study purposes, regions of interest from the same PEG blocks as in the prior protocol can be sectioned at 100 µm. In this instance, visualization of the cortical capillary network in and around microinfarcts is best achieved using immunoreactions against collagen IV (Figure 1 c-f). The primary antibody used below displays robust immunoreactivity in archival fixed brain tissue stored in formaldehyde solutions and achieves optimal penetration of free-floating sections at all optical planes during low-power microscopy. Recently, collagen IV immunohistochemistry was successfully applied to 40 µm paraffin sections by Decker et al.8 Here, we combined the collagen IV immunoreaction with aldehyde fuchsine staining (with or without Darrow red) or in combination with staining of vessel wall components vessel wall in 100 µm PEG sections (Figure 1 c,d).

1. Perform steps 1-3 as above.

2. Rinse sections briefly in deionized water, then reduce background immunoreaction by placing them for 30 min in a mixture of 10% methanol (100%) plus 10% hydrogen peroxide (30%) and 80 mL Tris. Rinse 5 min in Tris or PBS buffer. Agitate gently on a shaking table.

3. Perform blocking with bovine serum albumin solution for 1 h on a shaking table (gentle setting). Rinse sections twice in 70% ethanol to remove excess dye. When only aldehyde fuchsine staining is desirable, proceed to steps 5 and 6 immediately below.

4. Alternatively, transfer sections to deionized water and place into Darrow red solution for 1 h or more (agitate gently on a shaking table). To do so, dissolve 0.25 g Darrow red (Aldrich/Merck, Darmstadt, Germany) in 1000 mL deionized water, add 15 mL glacial acetic acid, boil for 20 min, filter solution, and add 250 mL of 0.2 Mol (i.e., 6.9 g sodium acetate trihydrate in 250 mL deionized water) sodium acetate solution (Merck, Darmstadt, Germany).

5. Dehydrate sections through a graded ethanol series. To obtain flatness, place free-floating sections between sheets of filter paper and compress gently between porcelain desiccator plates (140 mm diameter, Inv #119c, Morgan Technical Ceramics Haldenwanger, Germany, or Inv #8104, Lehman Scientific, Wrightsville, PA, USA) in 70%, 96% (30 min each), and 100% ethanol (30 min).

6. Remove sections from filter paper and dehydrate for another 30 min in 100% ethanol. A thorough dehydration step is crucial when using 300 µm sections. Clear in xylene twice (30 min each), coverslip in a high-refractive medium or synthetic resin (Histomount, National Diagnostics, Atlanta, GA, USA), and mount on slides.

Figure 1. Cortical microinfarcts in PEG-embedded sections of 300 µm and 100 µm thickness. The combination of aldehyde fuchsine and Darrow red in a 300 µm section (framed area in panel a is shown in greater detail in panel b) permits easy recognition of another cortical microinfarct (same individual as in Figure 2, Brodmann area 4). a) Arrow at far-left points to an unstained blood vessel; asterisks indicate gray matter areas that have undergone neuronal loss and glial cell loss; large funnel-like accumulations of aldehyde fuchsine-positive macrophages are oriented radially through multiple cellular cortical layers of the primary motor neocortex. c-f) The combination of aldehyde fuchsine staining with an immunoreaction against collagen IV (brown chromogen 3,3’-diaminobenzidine tetrahydrochloride, DAB) and a counterstain (eosin-phosphotungstic acid-aniline blue, EPA) permits more detailed study of cortical microinfarctions (100 µm, 74-year-old male); framed areas in panels c and e are shown in greater detail in panels d and f; erythrocytes at upper left in panel d are stained for erythrosine. e,f) Aldehyde fuchsine staining combined with an immunoreaction for collagen IV and Darrow red displays thickening and puckering of the damaged capillary network in a presumably more recent microinfarction: in contrast to the situation in older microinfarcts (c,d), the central portion of the lesion in panels e and f contains only a few aldehyde fuchsine-positive macrophages. Scale bars: c,e) 200 µm; f) 100 µm. Objectives used: a) 2x; b,c,e) 4x; d,f) 10x.
albumin for 90 min to prevent non-specific binding, rinse 5 min in Tris or PBS buffer.
4. Incubate sections with primary polyclonal antibody against collagen IV (Abcam ab6586, Cambridge, UK; 1:5000) at room temperature for 18 h on a shaking table (gentle setting). Rinse 5 min in Tris or PBS buffer.
5. Transfer specimens to secondary biotinylated antibody (anti-rabbit IgG, BA-1000, Vector Lab, Burlingame, CA, USA, 1:200) at room temperature for 90 min on a shaking table (gentle setting). Rinse 5 min in Tris or PBS buffer.
6. Visualize immunoreaction with the avidin-biotin complex (ABC Elite, PK-6100, Vector Lab) at room temperature for 120 min. Rinse 5 min in Tris or PBS buffer.
7. Transfer sections into 3,3'-diaminobenzidine tetrahydrochloride (DAB, D5637 Sigma, Taufkirchen, Germany) for 30 s to 5 min, rinse thoroughly in deionized water.
8. Transfer sections to a solution of 0.1 g erythrosine (Merck, Darmstadt, Germany) in 100 mL deionized water for 10 min. Rinse 2 min in deionized water.
9. Dissolve 5 g phosphotungstic acid (Merck) in 100 mL 96% in ethanol. Agitate sections gently in phosphotungstic acid solution for 30 min or longer on a shaking table until excess erythrosine is removed except for that in erythrocytes. Rinse 2 min in deionized water.
10. Prepare aniline blue stock solution by dissolving 1 g phosphotungstic acid and 0.3 g aniline blue (Chroma Waldeck, Münster, Germany) in 100 mL deionized water. Then, add 1 mL glacial acetic acid. Stain specimens in 1 mL stock solution in 50 mL deionized water on a shaking table at gentle setting for 20-30 min. Rinse 1 min in 0.5% acetic acid and dehydrate sections as in protocol I, steps 5 and 6.
11. Addition of cinnamaldehyde (nD 1.6219 at 20°C) to the mounting medium [8 mL Histomount and 2 mL cinnamaldehyde (Merck-Schuchardt, Hohenbrunn, Germany)] increases the refractive index (nD). When cover-slipping, sections should be mounted in undiluted cinnamaldehyde after the second clearing in xylene.
12. Alternatively to steps 8-11, perform counterstaining of Nissl material by transferring sections into Darrow red solution for 1 h or more and agitate gently on a shaking table (follow steps 4-6 in protocol I).

Results and Discussion

In protocol I, microinfarcts are recognizable as funnel-like entities that contain radially-oriented accumulations of aldehyde fuchsin-positive macrophages (Figures 1a,b and 2b). Protocol II stains not only the microinfarcts but also the affected capillary network, at whose center accumulations of dead neurons and aldehyde fuchsin-positive macrophages cluster (Figure 1c). Very early puckering within the capillary network may indicate recent microinfarcts (Figure 1e) where very little proliferation of aldehyde fuchsin-positive macrophages was observed at the center of such lesions (Figure 1f) in contrast to the dense accumulations of macrophages seen in other infarctions (Figure 1c,d). Pigment-Nissl staining, which combines aldehyde fuchsin for selective demonstration of neuronal loss – extraneuronal remnants of lipofuscin indicate the positions of dead nerve cells – with a Nissl stain (e.g., Darrow red) for topographical overview in the human brain (Figure 2a), has been in use for many years.13,18,19 However, this stain has not been previously employed to visualize cortical microinfarcts. Here, aldehyde fuchsin effectively stained macrophages in human
brain tissue (Figure 2b) and, thus, it can be used instead of CD68 immunoreactions (Figure 3). Here, we also combined pigment-Nissl staining with a sensitive immunoreaction (collagen IV)\(^{16}\) that also renders possible study of cortical microinfarcts with high spatial resolution (Figure 1 c-f). A drawback of both protocols is that they were performed on unconventionally large and thick free-floating tissue sections (≥100 µm), which, although suitable for research purposes, are not amenable to the demands of routine histopathology laboratories that rely on standard embedding cassettes and automated stainers. For this reason, we previously adapted our tissue processing procedures to also make possible the use of 70-100 µm paraffin-embedded free-floating sections,\(^{15}\) a method that has been employed successfully elsewhere.\(^{20,21}\)

Current non-invasive \textit{in vivo} MRI protocols are constantly undergoing improvement to diagnose and treat patients with cerebrovascular changes. While having attained greater importance in the daily clinical routine, such protocols need interpretation. In particular, efforts are in progress to relate \textit{in vivo} MRI results directly to histopathological results won from post-mortem study of brains with cortical microinfarcts.\(^{2}\) Our staining protocols (Figures 1 and 2 a,b) represent an improvement of and an alternative to current methods in thin H&E as well as CD68 sections (Figures 2 c-e and 3 a-c), and we plan to apply them to compare \textit{in vivo} MRI datasets with neuropathology of cortical microinfarcts in corresponding 100-300 µm serial whole brain sections.

Figure 3. Paraffin sections (5 µm) comparing CD68 immunoreactions (a-c) with aldehyde fuchsin staining of macrophages (d,e) in the cortical microinfarct shown in Figure 2 b-e. a) The 5 µm section immediately adjacent to the H&E stained section shown in Figure 2c was immunostained for CD68-positive macrophages (here, without nuclear counterstaining); this technique readily recognizes the microinfarct and permits evaluation of the form and extent of the CD68-positive macrophages. b-e) The next adjacent 5 µm section (d,e) only underwent aldehyde fuchsin staining to visualize macrophages (again, without nuclear counterstaining); the framed area in panel d is displayed in greater detail in panel e. The macrophages in one and the same 5 µm section are thus compared as they appear in aldehyde fuchsin staining (d, e) and CD68 immunoreaction (b, c), thereby making it possible to use aldehyde fuchsin staining in standard paraffin or thick tissue sections instead of CD68 immunohistochemistry. Scale bars: d) 100 µm; e) 25 µm. Objectives used: a) 4:1; b,d) 10:1; c,e) 40:1.

References